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CLAIMS

1. A method of [¹¹C]-radiolabelling a phenothiazine compound or a phenothiazinelike compound, wherein:

said compound has a polycyclic core of three six-membered rings fused together in a linear fashion and denoted the A-ring, B-ring, and C-ring, where the B-ring is the "middle" ring;

said polycyclic core is partially-aromatic or fully-aromatic;

said polycyclic core has 14 ring atoms, including exactly 1 or exactly 2 ring heteroatom(s), each of which is independently selected from N, O, and S;

the remainder of said ring atoms being C;

said exactly 1 or exactly 2 ring heteroatom(s) form part of the B-ring, but not part of the A-ring or C-ring, and so are located at one or both of the "central" positions denoted by a hash-mark (#) in the following depiction of the polycyclic core:

said compound has a pendant group covalently attached to a ring atom of said polycyclic core;

said pendant group is independently:

a primary amino group;

a cationic primary imino group;

a secondary amino group;

a cationic secondary imino group;

a primary imino group; or

a secondary imino group;

said method comprising the step of:

reacting said phenothiazine compound or a phenothiazine-like compound with [11C]methyl trifluoromethanesulfonate (CF₃SO₂O¹¹CH₃);

thereby converting said pendant group to a corresponding [¹¹C]methyllabelled pendant group, respectively:

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- a [11C]methyl-labelled secondary amino group;
- a [11C]methyl-labelled cationic secondary imino group;
- a [11C]methyl-labelled tertiary amino group;
- a [11C]methyl-labelled cationic tertiary imino group;
- a [11C]methyl-labelled secondary imino group; or
- a [11C]methyl-labelled cationic tertiary imino group;

to give a [¹¹C]-radiolabelled phenothiazine or phenothiazine-like compound.

- A method according to claim 1, wherein said polycyclic core has 14 ring atoms, including exactly 2 ring heteroatoms, each of which is independently selected from N, O, and S.
 - 3. A method according to claim 1, wherein said polycyclic core has 14 ring atoms, including exactly 2 ring heteroatoms: N and S:

- 4. A method according to any one of claims 1 to 3, wherein said polycyclic core is fully-aromatic.
- 5. A method according to any one of claims 1 to 4, wherein said pendant group is independently attached to a ring carbon atom of said polycyclic core.
- 6. A method according to any one of claims 1 to 4, wherein said pendant group is independently attached to a ring carbon atom of said A-ring or C-ring, but not of said B-ring.
 - 7. A method according to any one of claims 1 to 4, wherein said pendant group is independently attached at one of the "distal" positions of said A-ring or C-ring, which positions are denoted by asterisks (*).

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- 8. A method according to any one of claims 1 to 7, wherein said pendant group is independently:
 - a secondary amino group or
 - a cationic secondary imino group;
 - and said corresponding [11C]methyl-labelled pendant group, respectively, is:
 - a [11C]methyl-labelled tertiary amino group; or
 - a [11Clmethyl-labelled cationic tertiary imino group.
- 9. A method according to any one of claims 1 to 7, wherein said pendant group is10 independently selected from:
 - $-NH_2$, -NHR, $=N^{(+)}H_2$, $=N^{(+)}HR$, =NH, and =NR;
 - wherein R is independently selected from C_{1-6} alkyl, C_{1-6} alkenyl, C_{1-6} alkynyl, C_{1-6} cycloalkyl, and C_{1-6} cycloalkenyl, and is optionally substituted with one or more groups selected from halo (e.g., fluoro, chloro, bromo, iodo), hydroxy, and C_{1-4} alkoxy;
 - and said corresponding [¹¹C]methyl-labelled pendant group, respectively, is:
 - $-NH-(^{11}CH_3),\ -NR-(^{11}CH_3),\ =N^{(+)}H-(^{11}CH_3),\ =N^{(+)}R-(^{11}CH_3),\ or\ =N-(^{11}CH_3).$
- 20 10. A method according to any one of claims 1 to 7, wherein said pendant group is independently selected from: -NHR and = $N^{(+)}HR$;
 - wherein R is independently selected from C_{1-6} alkyl, C_{1-6} alkenyl, C_{1-6} alkenyl, C_{1-6} cycloalkyl, and C_{1-6} cycloalkenyl, and is optionally substituted with one or more groups selected from halo (e.g., fluoro, chloro, bromo, iodo), hydroxy, and C_{1-4} alkoxy;
 - and said corresponding [11 C]methyl-labelled pendant group, respectively, is: -NR-(11 CH₃) or =N $^{(+)}$ R-(11 CH₃).
 - 11. A method according to claim 9 or 10, wherein R is independently C₁₋₄alkyl.
 - 12. A method according to claim 9 or 10, wherein R is independently -Me or -Et.
 - 13. A method according to claim 9 or 10, wherein R is independently -Me.

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14. A method according any one of claims 1 to 13, wherein said compound has, in addition to said pendant group, one or more additional substituents selected from:

amino (-NH₂), methylamino (-NHMe), dimethylamino (-NMe₂), ethylamino (-NHEt), diethylamino (-NEt₂), imino (=NH), methylimino (=NMe), ethylimino (=NEt), methyl (-Me), ethyl (-Et), fluoro (-F), chloro (-Cl), bromo (-Br), iodo (-I), oxo (=O), hydroxy (-OH), carboxy (-COOH), and protonated and deprotonated forms thereof.

15. A method according to claim 1, wherein the phenothiazine or phenothiazine-like compound is a compound of the following formula:

wherein:

each of R^1 , R^2 , and R^3 is independently -H, C_{1-6} alkyl, C_{1-6} alkenyl, C_{1-6} alkynyl, C_{1-6} cycloalkyl, and C_{1-6} cycloalkenyl, and is optionally substituted with one or more groups selected from halo (e.g., fluoro, chloro, bromo, iodo), hydroxy, and C_{1-4} alkoxy; and

M⁻ is an anion.

- 16. A method according to claim 15, wherein -NHR¹ is independently -NHMe.
- 17. A method according to claim 15 or 16, wherein -NR²R³ is independently -NH₂.
- 18. A method according to claim 15 or 16, wherein -NR²R³ is independently -NHMe.
- 25 19. A method according to claim 15 or 16, wherein -NR²R³ is independently -NMe₂.
 - 20. A method according to any one of claims 15 to 19, wherein M⁻ is independently a halide ion.
- 30 21. A method according to any one of claims 15 to 19, wherein M⁻ is independently Cl⁻.

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22. A method according to claim 1, wherein the phenothiazine or phenothiazine-like compound is Azure B:

and said [11C]-radiolabelled phenothiazine or phenothiazine-like compound is [N-methyl-11C]methylene blue:

- 23. A method according to any one of claims 1 to 22, wherein said reaction is performed in the presence of a Bronsted base.
- 24. A method according to any one of claims 1 to 22, wherein said reaction is performed in the presence of an alkali metal carbonate or bicarbonate.
- 25. A method according to any one of claims 1 to 22, wherein said reaction is performed in the presence of potassium carbonate (K₂CO₃).
 - 26. A method according to any one of claims 1 to 25, wherein said reaction is carried out in aqueous media.
- 27. A method according to any one of claims 1 to 25, wherein said reaction is carried out by introducing said [¹¹C]methyl trifluoromethanesulfonate into an aqueous solution or suspension of said phenothiazine or phenothiazine-like compound, to form a reaction mixture.
- 25 28. A method according to claim 27, wherein said aqueous solution or suspension further comprises a Bronsted base.

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| 29. | A method according to claim 27, wherein said aqueous solution or suspension further comprises an alkali metal carbonate or bicarbonate. |
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| 30. | A method according to claim 27, wherein said aqueous solution or suspension further comprises potassium carbonate (K_2CO_3). |
| 31. | A method according to any one of claims 27 to 30, wherein said reaction mixture is mixed for a mixing time of 1-30 minutes. |
| 32. | A method according to any one of claims 27 to 30, wherein said reaction mixture is mixed for a mixing time of 1-10 minutes. |
| 33. | A method according to any one of claims 27 to 32, wherein said reaction is carried out at 20°C-25°C. |
| 34. | A method according to any one of claims 27 to 32, wherein said reaction is carried out under an inert atmosphere. |
| 35. | A method according to any one of claims 27 to 32, wherein said reaction is carried out under argon. |
| 36. | A method according to any one of claims 1 to 35, further comprising the subsequent step of: purifying said [¹¹C]-radiolabelled phenothiazine or phenothiazine-like compound. |
| 37. | A method according to any one of claims 1 to 35, further comprising the subsequent step of: purifying said [11C]-radiolabelled phenothiazine or phenothiazine-like compound using ion exchange methods. |

38. A method according to any one of claims 1 to 35, further comprising the subsequent step of:

purifying said [¹¹C]-radiolabelled phenothiazine or phenothiazine-like compound using cation exchange methods.

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- 39. A method according to any one of claims 1 to 38, wherein the reaction and optional purification is performed in less than 60 minutes.
- 40. A method according to any one of claims 1 to 38, wherein the reaction and optional purification is performed in less than 45 minutes.
 - 41. A method according to any one of claims 1 to 38, wherein the reaction and optional purification is performed in less than 40 minutes.
- 10 42. A method according to any one of claims 1 to 41, which provides a radiochemical purity greater than 90%.
 - 43. A method according to any one of claims 1 to 42, which provides a radiochemical yield of at least 2%.
 - 44. A method according to any one of claims 1 to 43, which provides a specific average activity of at least 0.5 GBq/µmol.
- 45. A method according to any one of claims 1 to 44, which is partially or fully automated.
 - 46. A [11C]-radiolabelled phenothiazine or phenothiazine-like compound which is obtained by a method as defined in any one of claims 1 to 45.
- 25 47. A composition comprising a compound according to claim 46.

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- 48. A composition comprising a compound according to claim 46 and a pharmaceutically acceptable carrier or excipient.
- 30 49. A method of PET imaging which employs a compound according to claim 46.
 - 50. A method of PET imaging comprising the steps of:
 - (i) preparing a [¹¹C]-radiolabelled phenothiazine or phenothiazine-like compound using a method according to any one of claims 1 to 45;
 - (ii) introducing said compound into a subject; and
 - (iii) PET imaging (e.g., a part of, the whole of) the subject.

- 51. A compound according to claim 46 for use in a method of treatment of the human or animal body by therapy.
- 5 52. A compound according to claim 46 for use in a diagnostic or prognostic method practiced on the human or animal body.
 - 53. Use of a compound according to claim 46 in the manufacture of a medicament for use in the treatment of skin cancer (e.g., melanoma) or a tauopathy (e.g., Alzheimer's disease).
 - 54. Use of a compound according to claim 46 in the manufacture of a medicament (e.g., a diagnostic or prognostic reagent) for use in the diagnosis or prognosis of skin cancer (e.g., melanoma) or a tauopathy (e.g., Alzheimer's disease).

55. Use of:

 (i) a phenothiazine compound or a phenothiazine-like compound, wherein: said compound has a polycyclic core of three six-membered rings fused together in a linear fashion and denoted the A-ring, B-ring, and C-ring, where the B-ring is the "middle" ring;

said polycyclic core is partially-aromatic or fully-aromatic;

said polycyclic core has 14 ring atoms, including exactly 1 or exactly 2 ring heteroatom(s), each of which is independently selected from N, O, and S;

the remainder of said ring atoms being C;

said exactly 1 or exactly 2 ring heteroatom(s) form part of the B-ring, but not part of the A-ring or C-ring, and so are located at one or both of the "central" positions denoted by a hash-mark (#) in the following depiction of the polycyclic core:

said compound has a pendant group covalently attached to a ring atom of said polycyclic core;

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said pendant group is independently:

- a primary amino group;
- a cationic primary imino group;
- a secondary amino group;
- a cationic secondary imino group;
- a primary imino group; or
- a secondary imino group;

and

(ii) [11 C]methyl trifluoromethanesulfonate (CF $_3$ SO $_2$ O 11 CH $_3$); in the manufacture of a medicament for use in the treatment of skin cancer (e.g., melanoma) or a tauopathy (e.g., Alzheimer's disease).

56. Use of:

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(i) a phenothiazine compound or a phenothiazine-like compound, wherein: said compound has a polycyclic core of three six-membered rings fused together in a linear fashion and denoted the A-ring, B-ring, and C-ring, where the B-ring is the "middle" ring;

said polycyclic core is partially-aromatic or fully-aromatic;

said polycyclic core has 14 ring atoms, including exactly 1 or exactly 2 ring heteroatom(s), each of which is independently selected from N, O, and S;

the remainder of said ring atoms being C;

said exactly 1 or exactly 2 ring heteroatom(s) form part of the B-ring, but not part of the A-ring or C-ring, and so are located at one or both of the "central" positions denoted by a hash-mark (#) in the following depiction of the polycyclic core:

said compound has a pendant group covalently attached to a ring atom of said polycyclic core;

said pendant group is independently:

a primary amino group;

a cationic primary imino group;

a secondary amino group;

a cationic secondary imino group;

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a primary imino group; or a secondary imino group;

and

- (ii) [¹¹C]methyl trifluoromethanesulfonate (CF₃SO₂O¹¹CH₃); in the manufacture of a medicament (e.g., a diagnostic or prognostic reagent) for use in the diagnosis or prognosis of skin cancer (e.g., melanoma) or a tauopathy (e.g., Alzheimer's disease).
- 57. A method of manufacturing a medicament for use in the treatment of skin cancer (e.g., melanoma) or a tauopathy (e.g., Alzheimer's disease) which includes the steps of a method according to any one of claims 1 to 45.
 - 58. A method of manufacturing a medicament for use in the diagnosis or prognosis (e.g., of skin cancer (e.g., melanoma) or a tauopathy (e.g., Alzheimer's disease) which includes the steps of a method according to any one of claims 1 to 45.
 - 59. A method of treatment of skin cancer (e.g., melanoma) or a tauopathy (e.g., Alzheimer's disease) in a patient, comprising administering to said patient a therapeutically-effective amount of a compound according to claim 46.
 - 60. A method of treatment of skin cancer (e.g., melanoma) or a tauopathy (e.g., Alzheimer's disease) in a patient, comprising the steps of:
 - (i) preparing a [¹¹C]-radiolabelled phenothiazine or phenothiazine-like compound using a method according to any one of claims 1 to 45;
 - (ii) administering to said patient a therapeutically-effective amount of said [¹¹C]-radiolabelled phenothiazine or phenothiazine-like compound.
- 61. A method of diagnosis or prognosis of skin cancer (e.g., melanoma) or a tauopathy (e.g., Alzheimer's disease) which employs a compound according to claim 46.

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- 62. A method of diagnosis or prognosis of skin cancer (e.g., melanoma) or a tauopathy (e.g., Alzheimer's disease) comprising the steps of:
 - (i) preparing a [¹¹C]-radiolabelled phenothiazine or phenothiazine-like compound using a method according to any one of claims 1 to 45;
 - (ii) introducing said [¹¹C]-radiolabelled phenothiazine or phenothiazine-like compound into the subject;
 - (ii) determining the presence and/or location and/or amount of [¹¹C]-radiolabelled phenothiazine or phenothiazine-like compound in (e.g., a part of, the whole of) the subject;
 - (iii) correlating the result of the determination made in (ii) with a disease condition of the subject.

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